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Fabrication of Bi₂O₂CO₃/g-C₃N₄ heterojunctions for efficiently photocatalytic NO in air removal: *In-situ* self-sacrificial synthesis, characterizations and mechanistic study



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ABSTRACT

Layer-structured Bi₂O₂CO₃/g-C₃N₄ heterojunction photocatalysts were successfully prepared via onepot hydrothermal method for the first time, in which graphitic carbon nitride $(g-C_3N_4)$ served as the self-sacrificial reagent to supply carbonate anions simultaneously. Our results showed that the in situ fabricated Bi₂O₂CO₃/g-C₃N₄ heterojunction exhibited superior visible-light-driven photocatalytic activity for NO photocatalytic oxidation, which can be ascribed to the morphology and structure modulation during the sacrificial synthesis processes. Heterojunctions formation pathways underlying temperatureand time-dependent structure evolution were discussed in detail. The sample fabricated at 160 °C for 12 h (BOC-CN-160) showed high stability and durability, and the highest NO removal rate which is up to 34.8% under visible light irradiation. Results from photocurrent tests and electrochemical impedance spectroscopy (EIS) demonstrated that the BOC-CN-160 sample presents much more effective interface charge separation efficiency, which can contribute to its remarkably improved photocatalytic performance. Reactive radicals during the photocatalysis processes were identified by electron spin resonance (ESR) study. Combined with the quantification of reaction intermediates, the photocatalytic degradation mechanism of NO over Bi₂O₂CO₃/g-C₃N₄ heterojunction photocatalyst was proposed. The novel approach developed in this study may be further extended to synthesize a series of novel and highly efficient g-C₃N₄-based carbonate heterojunction photocatalysts for visible light-harvesting and energy conversion applications.

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1. Introduction

With the rapid urbanization and development of China, the concentration levels of PM $_{2.5}$ (PM $\leq 2.5~\mu m$ in aerodynamic diameter) in major megacities remains remarkably higher than that of the National Ambient Air Quality Standard [1–3]. Previous studies showed that secondary organic aerosols (SOAs), which are

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formed via the photochemical reactions with NO_x (NO+NO₂=NO_x) and volatile organic compounds (VOCs) as precursors, contributed significantly to the mass concentrations of PM_{2.5} [4–6]. As an alternative to conventional approaches (selective catalytic reduction and wet scrubbing, etc.) with high cost adopted for high-concentration NO_x treatment [7], semiconductor photocatalysis has gained considerable attention not only because of its potential value in addressing the worldwide energy shortage, but also because of its eco-friendly merit and high efficiency for the remediation of NO_x at part-per-billion (ppb) levels [8–13].

A suitable band gap structure of semiconductor photocatalyst is essential for light harvesting and photo-induced carriers separating and migrating, which facilitates the occurring of redox reaction thermodynamically [14]. Single semiconductor cannot fulfill the

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above harsh terms simultaneously. Hence, numerous strategies, such as element doping [15], noble deposition [16], and surface modification [17], have been developed to improve the photocatalytic activities of catalysts in the past decade. Recently, exploring semiconductor/semiconductor photocatalytic heterojunctions that mainly contain type-I/II/III heterojunctions, p/n-n heterojunctions, Z-scheme system, and homojunction for photocatalytic performance improvement has been paid more and more attention [18,19]. In contrast to single semiconductors, heterostructure composites not only effectively restrain the recombination of photoinduced e⁻-h⁺ pairs, but also markedly endow nanohybrids with enhanced properties or unique features through their synergistic catalytic effects [20].

 $Bi_2O_2CO_3$, consisted of $[Bi_2O_2]^{2+}$ slabs interleaved between two slabs of CO₃²⁻, possesses an Aurivillius-layered structure that exerts a low mammalian toxicity and displays a wide band gap (3.1–3.5 eV) [21,22] (Fig. S1). The compound has been used in medical treatment for a long time and as a photocatalyst in recent years [21,23]. However, Bi₂O₂CO₃ photocatalysts can only absorb UV light (less than 5% of the solar light) and exhibit a poor charge carriers separation ability, which seriously restrict the photocatalytic performance of this type photocatalysts. Therefore, various methods have been recently employed to overcome the intrinsic limitations of Bi₂O₂CO₃ photocatalysts, mainly including element doping (N, C, CO₃²⁻, Bi, Ag-doping Bi₂O₂CO₃) [24-28], and heterojunction fabrication $(Bi_2O_2CO_3/BiVO_3, Bi_2O_2CO_3/BiOX (X=Cl, Br, and I), Bi_2O_2CO_3/\alpha$ Bi_2O_3 , $Bi_2O_2CO_3/Ag_2O$, and $Bi_2S_3/Bi_2O_3/Bi_2O_2CO_3$) [22,29-35]. Among these approaches, the fabrication of heterojunctions was preferred by many researchers because it can maximize the advantages of each component to offset their individual weaknesses. Graphitic carbon nitride (g-C₃N₄) is a metal-free, visible-light, covalent binary CN polymer organic semiconductor photocatalyst with a 2D layered structure which is analogous to that of graphite [36,37]. This material has attracted considerable interest due to its unique physicochemical property and electronic band structure [38], which has been potentially applied in fields that are related to solar energy conversion [39,40], environmental remediation [41,42], bioimaging [43], and material science [44,45].

Considering the unique features of Bi₂O₂CO₃ and g-C₃N₄, a rational strategy is to combine the merits of each component to fabricate heterojunction nanocomposites with intrinsic layerstructure. Various methodologies have been adopted to construct Bi₂O₂CO₃/g-C₃N₄ heterojunction in order to extend the optical absorption and improve the quantum yield of Bi₂O₂CO₃. For example, Zhang et al. [46] prepared (BiO)₂CO₃/g-C₃N₄ heterojunction photocatalysts by the deposition of (BiO)₂CO₃ onto the surface of g-C₃N₄ for the efficient capture of atmospheric CO₂ at room temperature. Xiong et al. [47] prepared flower-like g-C₃N₄/Bi₂O₂CO₃ microspheres with a high adsorption ability via chemical precipitation method. Zhang et al. [48] designed novel g-C₃N₄/Bi₂O₂CO₃ nanojunction photocatalysts through self-assembly and chemical precipitation. However, all of these mentioned synthesis strategies for Bi₂O₂CO₃/g-C₃N₄ heterojunction fabrication require the addition of extra carbonate sources (such as CO₂ or Na₂CO₃), and normally require two main steps. The pristine component is prepared firstly and then deposited with another component to construct heterojunctions. These procedures are difficult to construct heterojunctions with uniform structures because of compatibility issues among different components. Therefore, constructing heterojunction structures with homogeneous distribution of components remains challenging.

In this study, layer-structured $Bi_2O_2CO_3/g$ - C_3N_4 heterojunction photocatalysts were successfully prepared with a one-pot hydrothermal strategy *via* self-sacrificing a part of g- C_3N_4 to supply CO_3^{2-} for $Bi_2O_2CO_3$ formation for the first time. The pre-

pared Bi₂O₂CO₃/g-C₃N₄ heterojunctions exhibit a considerably higher photocatalytic activity than any single phase on NO in air degradation under visible light irradiation. The mechanisms underlying structural evolution, NO removal, and photocatalytic activity enhancement are discussed through the combination of experimental results and theoretical analysis. This novel synthesis strategy can be probably extended to synthesize a series of highly efficient, visible light-driven g-C₃N₄-based carbonate heterojunction photocatalysts.

2. Experimental

2.1. Synthesis of $Bi_2O_2CO_3/g$ - C_3N_4 heterojunction samples

All chemicals were used without further purification. Bi₂O₂CO₃/g-C₃N₄ composites (BOC/CN for short) were synthesized using a hydrothermal approach. In a typical process, 2.0 g of Bi(NO₃)₃·5H₂O was dispersed in 40 mL of HNO₃ (0.02 M) and stirred for 30 min. Then, 0.9 g of g-C₃N₄ was added to the above solution. After 45 min of ultrasonic treatment, 0.625 M NaOH was added to adjust the pH to 8.0-8.5. The mixture was transferred into a 100 mL Teflon-lined autoclave and heated for 12 h at 140 °C, 160 °C, or 180 °C or heated at 160 °C for 4, 8, or 12 h. After cooling down to room temperature naturally, the products were collected by filtration, washed thoroughly with deionized water and absolute ethanol for three times, dried at 60 °C for 12 h, and then denoted as BOC-CN-140, BOC-CN-160, and BOC-CN-180, and BOC-CN-160-4, BOC-CN-160-8, and BOC-CN-160-12, respectively. For comparison, Na₂CO₃ was used instead of g-C₃N₄ to synthesize Bi₂O₂CO₃ in the same manner at 160 °C, and g-C₃N₄ used in this experiment was prepared according to the previously reported method [49] (Supplementary information).

2.2. Characterization methods

The crystal phase was analyzed via X-ray diffraction (XRD) with Cu K\alpha radiation at a scan rate of $0.05^{\circ}2\theta/s$ ($\lambda = 1.5406 \,\text{Å}$, $40 \,\text{kV}$, 40 mA, PANalytical X' Pert PRO X-ray diffractometer). Samples were embedded in KBr pellets and then subjected to Fouriertransform infrared spectroscopy (FT-IR) on an FT-IR absorption spectrometer (Magna-IR 750, Nicolet, USA). X-ray photoelectron spectroscopy (XPS) was conducted using an X-ray photoelectron spectrometer (Thermo ESCALAB 250, USA). All binding energies were calibrated to the C 1 s peak at 284.7 eV of the surface adventitious carbon. Elemental analyses were carried out on a Elementar Vario EL instrument (Vario EL III, German, detection limit: 0.015%, standard deviation: ≤0.1% abs) with He purging for 20 s prior to testing. Scanning electron microscopy (SEM, JEOL JSM-6490, Japan) was used to characterize the morphology and elemental distribution maps of the obtained products. Transmission electron microscopy (TEM) was performed using a JEOL JEM-2100HR CM-120 (Japan) transmission electron microscope. Moreover, a nitrogen adsorption apparatus (ASAP 2020, USA) was used to obtain the nitrogen adsorption-desorption isotherms of the samples that were degassed at 150 °C prior to analysis. A Varian Cary 100 Scan UV-vis system equipped with a labsphere diffuse reflectance accessory was used to obtain the reflectance spectra of the catalysts over a range of 200-800 nm. Labsphere USRS-99-010 was employed as a reflectance standard. The intermediate and final products, nitrate and nitrite ions, were extracted by immersing the powders (about 0.05 g) into 6 mL deionized water and measured by using a Dionex-600 Ion Chromatograph (IC, Dionex Inc., Sunny-vale, CA, USA) after the photocatalytic activity test equipped with an IonPac AS14A column. The mobile phase was composed of a mixture of 1.8 mM Na₂CO₃ and 1.7 mM NaHCO₃ at a flow rate of 1.20 mL/min, the volume of samples was $20~\mu L$. The detection limits for NO_2^- and NO_3^- are $0.0050~\mu g~m L^{-1}$ and $0.0256~\mu g~m L^{-1}$. Samples for electron spin-resonance spectroscopy (ESR, ER200-SRC, Bruker, Germany) were prepared by mixing 0.05~g of the as-prepared photocatalyst in a 25~mM 5.5'-dimethyl-1-pirroline-N-oxide (DMPO) solution with a 50~mL aqueous dispersion for DMPO- $^{\bullet}$ OH or a 50~mL alcohol dispersion for DMPO- $^{\bullet}$ O $_2^-$ under irradiation with 420~mm visible light.

2.3. Photoelectrochemical measurements

Photoelectrochemical properties of the as-prepared samples were evaluated using a Parstat 4000 electrochemical workstation (USA) in a conventional three-electrode cell, with a platinum plate and an Ag/AgCl electrode as the counter and reference electrodes, respectively. The photocurrent-time curves were measured at 0.2 V versus Ag/AgCl in 0.5 mol/L Na₂SO₃ at ambient temperature under a 300 W Xe lamp (microsolar 300, Perfectlight, China) with 420 nm. Electrochemical impedance spectroscopy (EIS) was measured at a frequency range of 0.1 Hz to 100 kHz with an 5 mV voltage amplitude under an open-circuit voltage in a 1 mmol/L K₃Fe(CN)₆ and $\rm K_4Fe(CN)_6$ solution.

2.4. Photocatalytic activity test

The photocatalytic activity of the as-prepared samples were investigated in terms of NO removal at ppb levels in a continuous flow reactor system (Fig. S2) at ambient temperature. The rectangular reactor, which was fabricated from stainless steel and covered with quartz glass, possessed a volume capacity of $4.5 L (30 cm \times 15 cm \times 10 cm)$. A 300 W commercial Xe arc lamp (microsolar 300, Perfectlight, China) was vertically placed outside the reactor. UV light in the light beam was removed by adopting a UV cutoff filter (420 nm). The photocatalyst (0.1 g) was coated onto one dish (10.0 cm in diameter), which was subsequently pretreated at 60 °C to remove water in the suspension. NO gas was supplied by a compressed gas cylinder at a concentration of 48 ppm (N₂ balance) in accordance with the standard of the National Institute of Standards and Technology. The initial NO concentration was diluted to 400 ppb by an air stream supplied by a zero-air generator (Model 1001, Sabio Instruments LLC, Georgetown, TX, USA), and the flow rate was controlled at 3 L/min. The desired relative humidity level of NO flow was controlled at 70% by passing zero air streams through a humidification chamber. The lamp was turned on when the adsorption-desorption equilibrium was achieved. The concentration of NO was continuously measured using a chemiluminescence NO analyzer (Model 42c, Thermo Environmental Instruments Inc., Franklin, MA, USA). This apparatus was used to monitor NO and NO2 concentrations at a sampling rate of $0.6 \, L/min$. The removal ratio (η) of NO was calculated as η (%)=(1-C/C₀) × 100, where C and C₀ are the concentrations of NO in the outlet and feeding streams, respectively.

3. Results and discussion

3.1. Phase structure and chemical compositions

Fig. 1a shows the XRD patterns of the as-prepared BOC, BOC-CN-160, and CN samples. Apparently, all diffraction peaks for BOC can be readily indexed to pure tetragonal $Bi_2O_2CO_3$ (a=3.865 Å, b=3.865 Å, c=13.675 Å, JCPDS no. 41-1488) without other diffraction peaks being detected [50]. Two distinct diffraction peaks located at $2\theta=13.1^{\circ}$ and 27.6° in g-C₃N₄ are attributed to the (100) crystal plane of tri-s-triazine units and the (002) crystal plane of interlayer stacking of aromatic segments, respectively [38,51]. The typical diffraction peaks of BOC and CN can also be observed in the

BOC-CN-160 composite, demonstrating that Bi₂O₂CO₃/g-C₃N₄ heterojunction has been successfully constructed by self-sacrificing parts of g-C₃N₄ as CO₃²⁻ provider. It is interesting to find that the (002), (004), and (006) peaks corresponding to the {001} facets of BOC exhibited overwhelmingly higher relative intensity than those of the heterojunction. Moreover, the intensity of (002) peaks belonging to CN decreases significantly because of the selfsacrifice reaction. It could be deduced that the BOC {001} facet growing in situ from the CN {002} facet leads to the simultaneous decrease in intensity of the (002) peaks in CN and the (002), (004), and (006) peaks in BOC. These results imply that the coupling of CN-BOC heterojunctions probably occurs between the BOC {001} and CN {002} facets. In addition, the phase structure of the obtained products evolved obviously as revealed by the results from temperature-dependent experiments. As shown in Fig. S3, the signal for Bi₂O₂CO₃ phase cannot be detected, and the main diffraction peaks for BOC-CN-140 can be indexed to tetragonal bismuth oxide nitrate hydroxide (Bi₂O₂(OH)NO₃, Fig. S4 presents the schematic crystal structure of Bi₂O₂(OH)NO₃), indicating that anions (NO₃⁻ and OH⁻) cannot overcome the kinetic barrier to exchange with CO₃²⁻ under this hydrothermal condition, and [Bi₂O₂]²⁺ covering the CN surface leads to the weak intensity of the CN feature peaks in BOC-CN-140. After hydrothermal treatment at 180 °C, a well-crystallized tetragonal Bi₂O₂CO₃ is formed, and no obvious CN peaks can be observed, which implies g-C₃N₄ may be decomposed almost completely in aqueous solution reaction under this condition

The FT-IR spectra of the BOC, BOC-CN-160, and CN are shown in Fig. 1b. The following typical bands of pure BOC are observed: an absorption band at 845 cm⁻¹, which is assigned to the out-ofplane bending mode of CO₃²⁻, and an absorption band at 1390 and 1467 cm⁻¹, which is assigned to the antisymmetric vibration mode of CO_3^{2-} [52]. For pure CN, the intensive bands in the 1200-1700 cm⁻¹ region indicate the stretching vibration of the heptazine heterocyclic ring (C_6N_7) units. The peak at $807 \, \text{cm}^{-1}$ is associated with the breathing mode of the heptazine ring system, whereas the absorption band at 890 cm⁻¹ is assigned to the deformation mode of N-H. In addition, the broad band at $3400-3600 \,\mathrm{cm}^{-1}$ is due to physically adsorbed H₂O [42,53]. The BOC-CN-140 sample shows the characteristic absorption bands of both Bi₂O₂(OH)NO₃ [54] and g-C₃N₄ (Fig. S5). The BOC-CN-160 sample shows the characteristic absorption bands of both BOC and g-C₃N₄, thereby confirming the co-presence of BOC and g-C₃N₄ components. The results of XRD and FT-IR indicate that Bi₂O₂CO₃/g-C₃N₄ heterojunction can be successfully fabricated by this in situ method using g-C₃N₄ as the sacrificial agent. In BOC-CN-180 sample (Fig. S5), the characteristic absorption bands of BOC were quite stronger than CN, which means little parts of undecomposed CN existed in BOC-CN-180.

The surface composition and coupling mode among atoms in the CN-BOC-160 heterojunctions are further analyzed by XPS. The surveyed spectra in Fig. 2a implies the presence of Bi, O, C, and N in the prepared nanocomposite sample without other element signals being detected. The two peaks at 164.2 and 158.9 eV in BOC are associated with Bi $4f_{5/2}$ and Bi $4f_{7/2}$ of Bi³⁺ [55,56], respectively (Fig. 2b). The distinct binding energy drifts observed in BOC-CN-160 can be attributed to the change in inner electron density caused by the formation of the BOC-CN heterojunction. In the C 1s region (Fig. 2c), the sharp peak around 284.6 eV is attributed to the pure graphitic species in the CN matrix [51], and the peak at 284.7 eV in BOC is assigned to adventitious carbon species from XPS measurement. The C 1s peak around 288.0 eV in the BOC-CN-160 sample can be assigned to the overlapping peaks of carbonate ion in BOC and the sp^2 -hybridized carbon in trizine rings (N–C=N) [51,57]. In the N 1s region (Fig. 2d), the main N 1s peak around the binding energy of 398.2 eV can be assigned to the sp^2 hybridized aromatic N

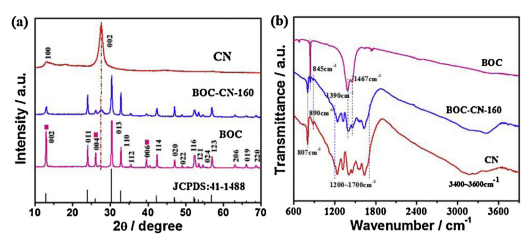
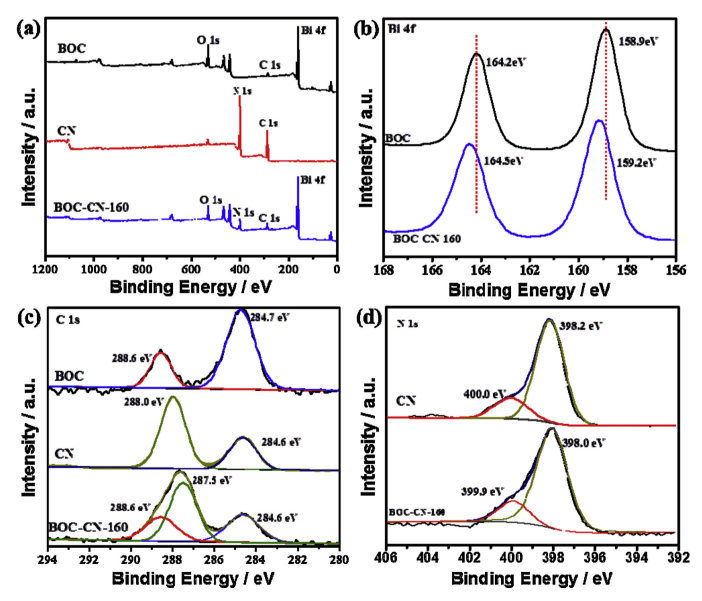


Fig. 1. XRD patterns (a) and FT-IR spectra (b) of pristine BOC, BOC-CN-160, and CN.



 $\textbf{Fig. 2.} \ \ \text{Survey XPS spectra (a) and high-resolution XPS spectra of Bi } \ 4f(b), C \ 1s(c), \text{ and N } 1s(d) \ \text{of the as-prepared samples}.$

bonded to carbon atoms (C=N-C), and the weak peak at 400.0 eV is attributed to the tertiary nitrogen $N-(C)_3$ groups or $H-N-(C)_2$ [58] in both CN and BOC-CN-160 samples.

In order to investigate the constituent content of BOC-CN-160 sample quantificationally, the C and N elements were chosen to estimate the $Bi_2O_2CO_3$ and C_3N_4 contents in heterojunction by ele-

mental analysis method and the results were listed in Table S1. From the results, the C contents in BOC, BOC-CN-160, and CN are 2.347%, 14.964% and 36.835%, respectively. The N contents in BOC-CN-160, and CN are 22.940%, and 61.977%. Therefore, the mass contents of $Bi_2O_2CO_3$ and $g-C_3N_4$ in the CN-BOC-160 composites are estimated to be 56.994% and 36.566%, respectively.

3.2. Morphological structure and texture property

The morphologies of the representative samples are examined by using SEM and TEM analysis. As displayed in Fig. 3a, the BOC products are composed of large-scale, round pill-like nanoplates. A close-up view of the nanoplates in Fig. 3b indicates that the nanoplates are ca. $100-250\,\mathrm{nm}$ thick. The quadrate $Bi_2O_2CO_3$ nanoplatelets anchor on the bulk $g-C_3N_4$ surface to be assembled into $Bi_2O_2CO_3/g-C_3N_4$ nanocomposites, as shown in Fig. 3c. The $Bi_2O_2CO_3$ nanoplates are ca. $100\,\mathrm{nm}$ to $1.5\,\mathrm{\mu m}$ in length and $40-80\,\mathrm{nm}$ in thickness (Fig. 3d). The elemental mapping images of the same low region (Figs. 3e–h) indicate that Bi, O, N, and C are homogeneously distributed on the obtained heterojunction and that the extra C on the background is attributed to the signal from conductive sellotape.

Transmission electron microscopy (TEM) is used to investigate the morphological structures of the samples, as shown in Fig. 4. As displayed in Fig. 4a and consistent with SEM results (Fig. 3a), the large-scale, irregular pill-like nanoplates of the BOC sample are further verified and observed. The low-magnification TEM image (Fig. 4b) shows that individual CN consists of a typical layer structure with different sizes and irregular shapes. The square Bi₂O₂CO₃ nanosheets attach on the g-C₃N₄ surface to assemble Bi₂O₂CO₃/g-C₃N₄ nanocomposites, as shown in Fig. 4c. The HRTEM image of the BOC square nanosheet in the BOC-CN-160 sample reveals a lattice spacing of approximately 0.275 nm, which can be ascribed to the (110) plane of tetragonal BOC (JCPDS 41–1488), as shown in Fig. 4d. The single-crystalline nature of BOC in the BOC-CN-160 sample is confirmed by a selected-area electron diffraction (SAED) pattern, which is assigned to the [001] zone-axis diffraction spots of tetragonal BOC [23] (Fig. 4e). Based on these results, it can be assumed that the {001} facet of BOC in the BOC-CN-160 sample is exposed. Moreover, combining with the results of SEM and TEM, it is interesting to find that the dimension and thickness of the layers of Bi₂O₂CO₃ (Fig. 3d) and C₃N₄ (Fig. 4c) in the BOC-CN-160 sample are reduced and become much thinner than those of Bi₂O₂CO₃ (Fig. 3b) synthesized through the conventional method with the addition of extra carbonate precursors and those of original g-C₃N₄ (Fig. 4b), which can be ascribed to the occurrence of the g-C₃N₄ self-sacrifice effect during the synthesis processes. The HR-TEM image of g-C₃N₄ is shown in Fig. S6a, and the similar morphology in BOC-CN-160 heterojunction and meanwhile the (110) plane of BOC (lattice spacing of 0.274 nm) can be observed in Fig. S6b, suggesting the intimate interfacial contact between BOC and g-C₃N₄.

The nitrogen adsorption-desorption isotherms are displayed in Fig. S7. The results show all samples are of type IV (BDDT classification) [59], suggesting the presence of mesopores. The specific surface areas are 4, 27, and 10 m²/g for BOC, BOC-CN-160, and CN, respectively. The large specific surface area for BOC-CN-160 was associated with the morphological transformation, parts of bulk g-C₃N₄ with low surface area self-sacrifice and quadrate Bi₂O₂CO₃ nanoplatelets with high surface area formation, which can be evidenced by SEM and TEM.

3.3. Formation mechanism

According to the phase structure and morphologies analysis, a schematic illustration of the fabrication of $Bi_2O_2CO_3/g-C_3N_4$ heterojunctions using $g-C_3N_4$ as the self-sacrificial reagent to sup-

ply carbonate anions is proposed (Scheme 1). Bi₂O₂(OH)NO₃ and Bi₂O₂CO₃ are composed of [Bi₂O₂]²⁺ layers sandwiched between two slabs of anions or groups, facilitating the potential phase transformation from Bi₂O₂(OH)NO₃ to Bi₂O₂CO₃ in theory [58]. The heterojunction fabrication strategy in this study involves the formation of a [Bi₂O₂]²⁺ cage structure template and the process of anion exchange. As shown in Scheme 1, CO₃²⁻ does not exchange with NO₃⁻ and OH⁻ in Bi₂O₂(OH)NO₃ to form Bi₂O₂CO₃ under 140 °C, which means a high hydrothermal energy is required to overcome the kinetic barrier for anion exchange because of the higher stability of Bi₂O₂CO₃ than that of Bi₂O₂(OH)NO₃. Moreover, g-C₃N₄ suffers from decomposition in liquid water at the proper temperature, similar to mesoporous silica SBA-15, and generates NH₄⁺ and CO₃²⁻ during hydrothermal treatment [60,61]. However, few people can take advantage of this property for composite fabrication (Process 1). When the temperature is increased to 160 °C, the tetragonal Bi₂O₂CO₃ is obtained (Process 2), as evidenced by the XRD results (Fig. S3a). These assumptions were also supported by the results from time-dependent hydrothermal experiments to investigate the growth and ripening processes of Bi₂O₂CO₃/g-C₃N₄ heterojunctions at 160 °C (Fig. S3b). Fig. S3b shows that the Bi₂O₂(OH)NO₃ phase is gradually transformed into Bi₂O₂CO₃ as the reaction time is prolonged from 4h to 12h at 160°C (Process 3). Therefore, combining the experimental results regarding to the evolution of morphology and structure and the schematic description is Scheme 1, the formation process of the heterostructured Bi₂O₂CO₃/g-C₃N₄ composite in this study can be described by the following equations (Eqs. (A.1)-(A.6)):

$$g\text{-}C_{3}N_{4}(part) \, + \, 6H_{2}O \, \rightarrow \, 3CO_{2} + 4NH_{3} \tag{A.1} \label{eq:A.1}$$

$$NH_3 + H_2O \rightarrow NH_4^+ + OH^-$$
 (A.2)

$$CO_2 + 2OH^- \rightarrow CO_3^{2-} + H_2O$$
 (A.3)

$$2Bi^{3+} + NO_3^- + 5OH^- \rightarrow Bi_2O_2(OH)NO_3 + 2H_2O$$
 (A.4)

$$Bi_2O_2(OH)NO_3 + CO_3^{2-} \rightarrow Bi_2O_2CO_3 + NO_3^{-} + OH^{-}$$
 (A.5)

$$Bi_2O_2CO_3 + g-C_3N_4(others) \rightarrow Bi_2O_2CO_3/g-C_3N_4heterojunction(A.6)$$

3.4. Band gap structure of Bi₂O₂CO₃/g-C₃N₄ heterojunctions

In general, the photocatalytic activity of the photocatalyst is closely related to its band structure. The UV–vis DRS was used to investigate the optical properties of the obtained samples, as shown in Fig. 5. For the pure BOC sample, the basal absorption edge occurs at a wavelength shorter than 360 nm, whereas the absorption intensity of g–C₃N₄ shows absorption edges at 480 nm (Fig. 5a). In comparison to BOC, the absorption band edge of BOC-CN-160 exhibits an apparent red shift with the introduction of CN. This shift is ascribed to the interaction between CN and BOC semiconductors in heterojunctions. The band energies ($E_{\rm g}$), which are estimated from the intercept of tangents to the plots of $(\alpha h \nu)^{1/2}$ versus photo energy [7] (Fig. 5b), are 2.43 and 3.28 eV for CN and BOC, respectively. The VB and CB potentials of a semiconductor material can be estimated using the following empirical equations [34]:

$$E_{VB} = \chi - E^e + 0.5E_g$$

$$E_{\rm CB} = E_{\rm VB} - E_{\rm g}$$

where E_{VB}/E_{CB} is the valence/conduction band edge potential, and χ is the Sanderson electronegativity of the semiconductor, which is defined as the geometric average of the absolute electronegativity of the constituent atoms. E^e is the energy of free electrons on the hydrogen scale (about 4.5 eV versus NHE), and E_g is the band gap energy of the semiconductor. The χ -values for BOC and g-C₃N₄ are approximately 6.54 and 4.73 eV [34,62], respectively. On the basis

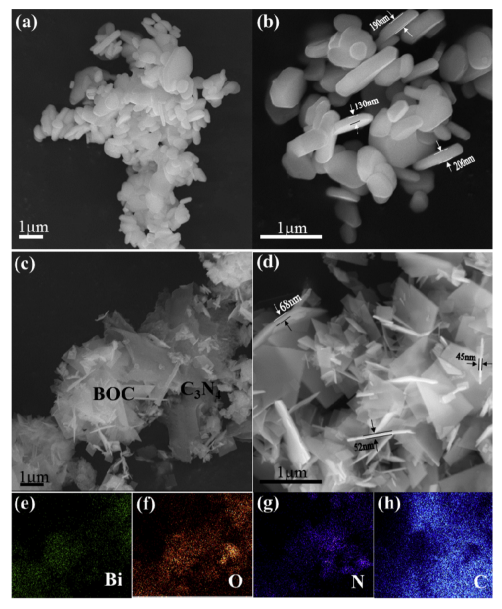


Fig. 3. SEM images of BOC at low (a) and high magnifications (b), BOC-CN-160 at low (c) and high magnifications (d), and elemental distribution maps of the same low region indicating the spatial distribution of Bi (green, e), O (red, f), N (purple, g), and C (blue, h). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

 Table 1

 Calculated CB, VB position, band gap energy, S_{BET} , NO removal ratios, and reaction rate constant for BOC, BOC-CN-160, and CN.

Samples	Calculated CB position (V)	Calculated VB position (V)	Band gap $E_{\rm g}$ (eV)	$S_{\rm BET}~({\rm m}^2/{\rm g})$	NO removal ratio η (%)	Reaction rate constant k (min ⁻¹)
ВОС	0.40	3.68	3.28	4	5.4	0.005
BOC-CN-160	_	_	_	27	34.8	0.077
CN	-0.99	1.44	2.43	10	21.6	0.050

of the above data, the top of the VB and the bottom of the CB of BOC are calculated to be 3.68 and 0.40 V, and the VB and CB of CN are estimated to be 1.44 and -0.99 V, respectively. The energy band structures of BOC and CN are listed in Table 1.

3.5. Photocatalytic activity and stability

The as-prepared BOC, BOC-CN-X (X = 140, 160, and 180), and CN samples were employed in the photocatalytic removal of NO under visible light irradiation in a continuous reactor to demon-

strate their potential capability in air purification. NO cannot be photolyzed under light irradiation without the aid of photocatalysis [25,63]. Figure 6a shows the variation of NO concentration over the as-prepared photocatalysts ($C/C_0\%$) with irradiation time. The NO concentrations decreased rapidly in the first 5 min because of the photocatalytic reaction and finally reach a stable value in all the cases. The removal ratios of NO by BOC, BOC-CN-180, BOC-CN-160, BOC-CN-140, and CN are 5.4%, 26.1%, 34.8%, 23.3%, and 21.6% after 30 min of visible light irradiation, respectively. In order to make a clearly quantitative comparison, we used the Langmuir-

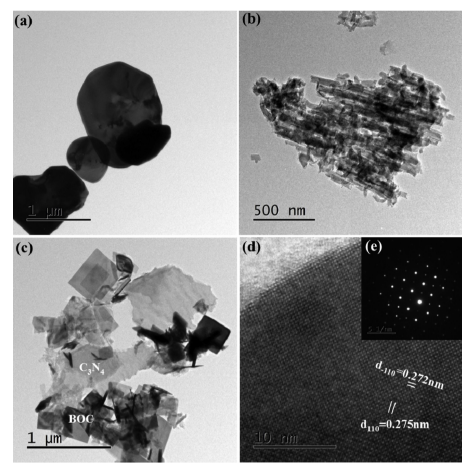
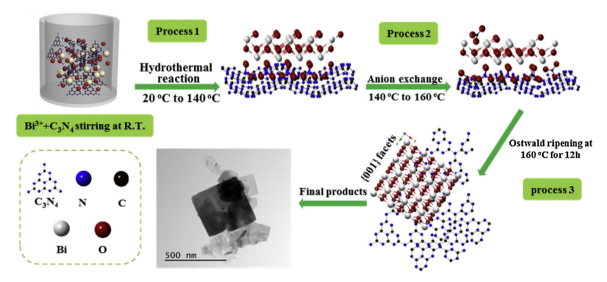


Fig. 4. Low-magnification TEM image of pristine BOC (a), CN (b), and BOC-CN-160 (c); HRTEM (d) and SAED images (e) of BOC square nanoplatelet of BOC-CN-160.



 $\textbf{Scheme 1.} \ \ \textbf{Scheme 2.} \ \ \textbf{Scheme 2.} \ \ \textbf{Scheme 3.} \ \ \textbf{Scheme 4.} \ \ \textbf{Scheme 4.} \ \ \textbf{Scheme 4.} \ \ \textbf{Scheme 5.} \ \ \textbf{Scheme 5.} \ \ \textbf{Scheme 6.} \ \ \textbf{Sc$

Hinshelwood model in reaction kinetics to describe the rates of NO photodegradation [64]. According to previous study [65], the initial photocatalytic degradation of NO was approximately simulated to follow mass-transfer-controlled pseudo-first-order rate reaction. (Details about Langmuir-Hinshelwood model and apparent reaction rate constant k calculations are given in the ESI†). The apparent reaction rate constant k of BOC-CN-160 (0.077 min⁻¹) is 15.4, 2.5, 1.6, and 1.5 times higher than those of BOC (0.005 min⁻¹),

BOC-CN-180 (0.031 min⁻¹), BOC-CN-140 (0.047 min⁻¹), and CN (0.050 min⁻¹), respectively (Fig. S8). Evidently, the BOC-CN-160 heterojunction exhibits the highest NO removal ratio and k among all the as-prepared samples. During the process of NO photocatalytic oxidation, NO₂, as a reaction intermediate, can form and hinder for the photocatalyst application because of its toxicity. Thus, the by-product of NO₂ is monitored online and selectivity is calculated (Fig. S9a and details about NO₂ selectivity are given in

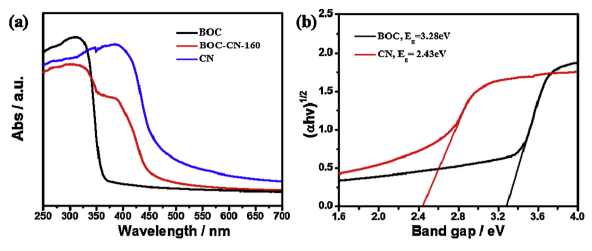


Fig. 5. UV-vis diffuse reflectance spectra (a) and plots of $(\alpha h \nu)^{1/2}$ versus the photo energy (b) of the as-synthesized BOC, BOC-CN, and CN nanocomposites.

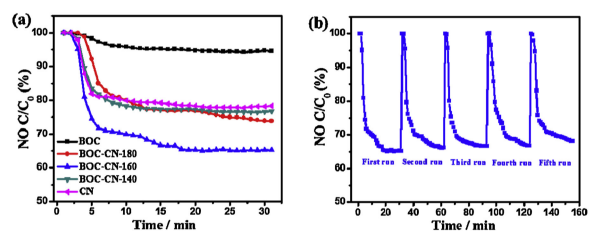


Fig. 6. (a) Visible light photocatalytic activities of the BOC, BOC-CN-*X* (*X* = 140, 160, 180), and CN samples for NO removal in air; (b) cycling runs for the photocatalytic degradation of NO over BOC-CN-160 under visible light irradiation.

the ESI†). The results show BOC-CN-160 displays the lowest conversion ratio of NO to NO₂ than others (Fig. S9b). Above results imply that the BOC-CN-160 heterojunction is more advantageous under realistic conditions because of outstanding NO degradation and NO₂ inhibition. The removal of NO can be attributed to the reaction between NO and photogenerated radicals, producing the final products of HNO₂ and HNO₃. In order to investigate the reaction pathways for the photocatalytic oxidation of NO_x, the accumulated NO₂⁻ and NO₃⁻ on the surface of BOC-CN-160 after single run reaction were extracted by deionized water and measured by ion chromatography (IC) method. It was found that the amounts of NO_3^- accumulated on BOC-CN-160 surface were 7.956 μ mol, but no NO2- was detected, which indicated that the oxidation of NO to NO₃⁻ was the major conversion process. It was found that the amount of NO degradation products as determined by IC analysis was lower than the total amount of removed NO (11.81 µmol) according to nitrogen mass balance. This can be attributed to the release of NO₂⁻ or NO₃⁻ into humidified air (RH 70%), or the generation of some new species undetectable by NO_x analyzer or IC, as Ai et al. reported [65]. Reproducibility and stability are important to photocatalysts for potential applications. To examine photocatalytic stability, cycling runs for the photocatalytic removal of NO with the BOC-CN-160 sample were performed under the same conditions. Fig. 6b illustrates the relationship between NO removal ratio and cycling times. After undergoing five repeated runs, the NO removal ratio remains high, indicating that BOC-CN-160 is stable in activity and can be used repeatedly. The used BOC-CN-160 sample is collected and analyzed by XRD and FT-IR (Fig. S10). Evidently, the crystal structure of used BOC-CN-160 sample is identical to that of the fresh sample, implying that the sample is stable and durable in structure.

3.6. Mechanism for NO removal and photocatalytic activity enhancement

To identify the formation of reactive radicals of the BOC-CN-160 sample accounting for NO removal, ESR spectroscopy with DMPO spin-trapping adducts was performed to allow the detection of superoxide radicals (DMPO-•O₂-) in methanol and hydroxyl radicals (DMPO-OH) in aqueous state. As shown in Fig. 7a, four apparent signals were generated, which can be attributed to •O2with BOC-CN-160 via the reduction of O2 with photo-generated electrons in methanol under visible light irradiation for 5 min. Meanwhile, no •O₂ - signal was observed in the dark under otherwise identical conditions. In Fig. 7b, DMPO-OH adduct signals that are weaker than DMPO-O₂ signals were also detected in the system containing the Bi₂O₂CO₃/g-C₃N₄ composite. The E_{VB} of g- C_3N_4 (1.44 V) is lower than the standard redox potential of $H_2O/^{\bullet}OH$ $(2.37\,V)$ but is higher than that of HNO_3/NO $(0.94\,V)$, indicating that the photogenerated holes of g-C₃N₄ could not oxidize H₂O to active species •OH but may oxidize NO (Eq. (B.2)) [66]. This result confirms that ${}^{\bullet}O_2^-$ is the dominant active species in NO_x oxidation. The ${}^{\bullet}OH$

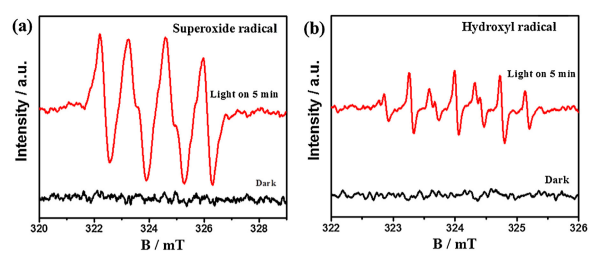


Fig. 7. DMPO spin-trapping ESR spectra of BOC-CN-160 in methanol dispersion for DMPO-*O2- (a) and in aqueous dispersion for DMPO-*OH (b).

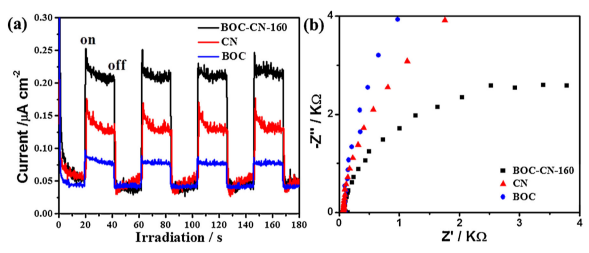


Fig. 8. Photocurrent transient (a) and Nyquist plots (b) for BOC-CN-160, CN, and BOC electrodes under visible light irradiation (λ > 420 nm).

radicals perform a minor function and should be generated *via* the $-{}^{\bullet}O_2^- \rightarrow {}^{\bullet}OH$ route (Eq. (B.4)) [42].

As revealed in Fig. 8a, transient photocurrent responses with good reproducibility were observed for the BOC-CN-160, CN, and BOC electrodes under visible light. These responses are directly correlated with the recombination efficiency of the photogenerated carriers. As forecasted, the weak photocurrent density of pure BOC can be detected because of the large energy gap, and decent photocurrent response singles of CN were detected. In contrast to both BOC and CN, the BOC-CN-160 nanohybrid exhibits a remarkably enhanced photocurrent density. This result confirms the more efficient charge separation of the BOC-CN-160 nanohybrid than BOC and CN because pure BOC acts as an electron acceptor in the heterojunction structure than that in the single phases. In addition, EIS measurements were carried out to further investigate the charge transfer resistance and separation efficiency of the charge carriers. Fig. 8b shows that BOC-CN-160 presents a smaller arc radius than the other samples under visible light irradiation ($\lambda > 420 \text{ nm}$). This result validates the high efficiency of charge transfer and separation of BOC-CN-160 [67]. In other words, the BOC-CN-160 nanohybrid exhibits lower resistance than the other samples and can facilitate the separation and immigration of photogenerated carriers under visible light irradiation.

Based on the above characterizations and analysis, the *in situ* fabricated $Bi_2O_2CO_3/g$ - C_3N_4 heterojunction exhibited superior visible-light-driven photocatalytic activity for NO photocatalytic

oxidation, which can be ascribed to the morphology and structure modulation during the sacrificial synthesis processes. The BOC-CN-160 heterojunction with the layered hierarchical structure of square BOC nanosheets show the highest BET surface area $(27 \,\mathrm{m}^2/\mathrm{g})$, which was favorable for reaction site increase, initial NO adsorption, and intermediates diffusion. More importantly, the synergistic effects of BOC-CN-160, including well-matched band structures, intimate interfacial contact, and effective separation of the photoinduced carriers, can be of great advantage to NO photocatalytic oxidation. The related mechanisms for NO removal and photocatalytic activity enhancement over Bi₂O₂CO₃/g-C₃N₄ heterojunction are proposed accordingly as illustrated in Fig. 9. g-C₃N₄ can be excited by visible light and generate photo-induced electrons and holes (Eq. (B.1)), but BOC cannot be subjected to identical conditions because of a wide band gap. The CB edge potential of g-C₃N₄ (-0.99V) is more negative than that of Bi₂O₂CO₃ $(0.40 \,\mathrm{V})$, allowing the excited electron on the g-C₃N₄ surface to transfer easily to Bi₂O₂CO₃ via the well-matched heterojunction under visible light irradiation. Thus, the photo-generated electrons and holes can be separated effectively, and the recombination of photogenerated charge would be sufficiently inhibited. Then, the photoelectrons electrons are trapped by O_2 in the surrounding air to form reactive ${}^{\bullet}O_2^-$ (Eq. (B.3)). NO_x reacts with the photo-generated -•O₂[−] and •OH radicals, and then NO₃[−] ions are produced (Eqs. (B.5)-(B.7)). The reaction processes of photocatalytic NO removal

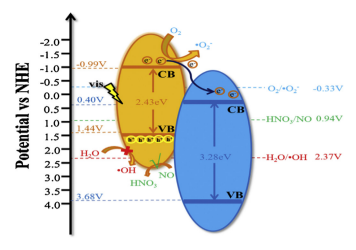


Fig. 9. Proposed photocatalytic mechanism for enhancing photocatalytic activity over Bi₂O₂CO₃/g-C₃N₄ heterojunctions under visible light irradiation.

by the $Bi_2O_2CO_3/g$ - C_3N_4 heterojunction are described as the following (Eqs. (B.1)-(B.7)):

$$g-C_3N_4 + hv \rightarrow e^- + h^+$$
 (B.1)

$$h^+ + NO + 2H_2O \rightarrow NO_3^- + 4H^+$$
 (B.2)

$$e^- + O_2 \rightarrow {}^{\bullet}O_2{}^- \tag{B.3}$$

$${}^{\bullet}\text{O}_{2}^{-} + 2\text{H}^{+} + 2\text{e}^{-} \rightarrow {}^{\bullet}\text{OH} + \text{OH}^{-}$$
 (B.4)

$$NO_X + {}^{\bullet}O_2^{-} \rightarrow NO_3^{-} \tag{B.5}$$

$$2^{\bullet}OH + NO \rightarrow NO_2 + H_2O \tag{B.6}$$

$$NO_2 + {}^{\bullet}OH \rightarrow NO_3^- + H^+ \tag{B.7}$$

4. Conclusion

Heterostructured Bi₂O₂CO₃/g-C₃N₄ nanocomposites with partial self-sacrificed g-C₃N₄ as the precursor of carbonate anion were successfully prepared by one-pot hydrothermal treatment for the first time. Compared with pristine Bi₂O₂CO₃ and g-C₃N₄, the BOC-CN-160 heterojunction sample exhibits dramatically enhanced visible light photocatalytic activity for the degradation of NO with the highest apparent reaction rate, attributing to favorable texture property and synergistic effects. In addition, the electrons, which are excited from g-C₃N₄ by irradiating BOC-CN-160 heterojunction under visible light, are trapped by O₂ in the surrounding air to form reactive ${}^{\bullet}O_2{}^-$ radicals, which were identified as the main active species in the photocatalytic reaction by ESR. The reaction mechanisms of photocatalytic NO removal by the Bi₂O₂CO₃/g-C₃N₄ heterojunction are discussed in detail. The in situ self-sacrificial synthesis strategy developed in this study may supply a new route for the fabrication of highly efficient C₃N₄-based carbonate heterostructures for environmental and energy applications in the future.

Notes

The authors declare no competing financial interest.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.apcatb.2016.06.027.

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